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Correspondence and
requests for materials
should be addressed to
J.D.B. (j.d.blount@
exeter.ac.uk) or K.E.P.
(kate.plummer@bto.
org)

* Current address:
British Trust for
Ornithology, The
Nunnery, Thetford,
Norfolk, IP24 2PU, UK.

Winter food provisioning reduces future breeding performance in a wild bird

K. E. Plummer^{1*}, S. Bearhop¹, D. I. Leech², D. E. Chamberlain³ & J. D. Blount¹

¹Centre for Ecology and Conservation, College of Life & Environmental Sciences, University of Exeter, Cornwall Campus, Penryn, Cornwall, TR10 9EZ, UK, ²British Trust for Ornithology, The Nunnery, Thetford, Norfolk, IP24 2PU, UK, ³Dipartimento di Scienze della Vita e Biologia dei Sistemi, Università degli Studi di Torino, Via Accademia Albertina 13, 10123 Turin, Italy.

Supplementation of food to wild birds occurs on an enormous scale worldwide, and is often cited as an exemplar of beneficial human-wildlife interaction. Recently it has been speculated that winter feeding could have negative consequences for future reproduction, for example by enabling low quality individuals to recruit into breeding populations. However, evidence that winter feeding has deleterious impacts on reproductive success is lacking. Here, in a landscape-scale study of blue tits (*Cyanistes caeruleus*) across multiple years, we show that winter food supplementation reduced breeding performance the following spring. Compared to unfed populations, winter-fed birds produced offspring that weighed less, were smaller, and had lower survival. This impairment was observed in parents that had received fat only, or in combination with vitamin E, suggesting some generality in the mechanism by which supplementary feeding affected reproduction. Our results highlight the potential for deleterious population-level consequences of winter food supplementation on wild birds.

Variation in food availability has profound effects on individual life-histories and the regulation of population dynamics¹. The supply of food to animal populations is often perturbed through human activities. In particular, supplementation of food to wild animals is extensively applied as a conservation tool and often yields almost immediate increases in productivity^{2–5}. However, studies of its downstream effects have highlighted some issues of concern such as biasing of primary sex ratios⁶, and increased disease transmission⁷. In addition to specific conservation projects, food supplementation to wildlife is in fact practiced on a far larger scale in gardens and backyards. For those species which use supplementary food in the UK, there is an estimated one feeder for every nine individual birds⁸, and similar levels of provisioning occur across much of the Western World^{9–12}. Supplementary feeding of birds therefore represents an enormous perturbation of natural resource availability.

Food supplementation is predicted to benefit wild birds¹¹, although a range of impacts seems possible. Recently it has been reported that food supplementation to birds whilst they are breeding can have immediate effects on diurnal activity patterns¹³ and result in reduced reproductive success¹⁴. However, it is unclear whether similarly deleterious effects may arise as a consequence of winter food supplementation. This is important because supplementary feeding is especially common during winter when there is an assumption that wild birds could benefit from being ‘helped’, yet there is considerable uncertainty about its impacts on future breeding performance^{10,15,16}. Carry-over effects occur where events or life history decisions in one season or year influence events in a subsequent season or year¹⁵. Such carry-over effects are thought to be widespread and to have important consequences for fitness in wild populations of animals, although there have so far been few experimental demonstrations of their existence¹⁵. Food supplementation in winter can have almost immediate benefits for survival (e.g.^{17,18}), and consequently it can result in increased recruitment and greater breeding densities the following spring¹⁹. However, only a single study to date has considered the effects of winter food supplementation on future reproductive success: winter feeding of blue tits (*Cyanistes caeruleus*) enabled birds to breed earlier and fledge more offspring during the following spring²⁰. In contrast, it has been speculated that winter food supplementation could also have deleterious impacts. For example it may result in receipt of an unbalanced diet, or lead to negative density dependent processes and changes to the phenotypic structure of breeding populations as a consequence of increased over-winter survival^{10,16,21}. Alternatively, food supplementation during winter could create an ‘ecological trap’ where birds receive inaccurate cues as to future natural food availability, and consequently make unsustainable investments in breeding¹⁶. If such effects are manifest they would be expected to cause reduced reproductive success at the population level.

Carry-over effects produced by winter supplementation might be influenced by the nutritional composition of foods. Items commonly provisioned to wild birds such as nuts, seeds and fats are rich in energy and lipophilic

Table 1 | Summaries of breeding parameters (mean \pm s.e.m) by winter feeding treatment and year between 2008–2010

Breeding parameter	Treatment	Year			Treatment mean
		2008	2009	2010	
Boxes occupied (%)	Unfed	37.7 \pm 5.9	38.9 \pm 11.8	47.2 \pm 8.2	41.3 \pm 4.7
	Fat	41.8 \pm 6.5	45.7 \pm 3.7	48.6 \pm 8.9	45.4 \pm 3.5
	Fat + Vit E	43.7 \pm 14.9	44.3 \pm 10.7	53.0 \pm 0.3	47.0 \pm 5.5
	Annual mean	41.1 \pm 5.1	43.0 \pm 4.8	49.6 \pm 3.6	
Lay date [1 = 1 April]	Unfed	27.6 \pm 1.2	24.3 \pm 0.9	29.6 \pm 0.9	27.3 \pm 0.6
	Fat	26.4 \pm 1.0	24.6 \pm 1.0	29.3 \pm 0.7	26.8 \pm 0.5
	Fat + Vit E	26.0 \pm 1.0	25.4 \pm 1.0	31.7 \pm 1.0	28.0 \pm 0.6
	Annual mean	26.6 \pm 0.6	24.8 \pm 0.6	30.2 \pm 0.5	
Clutch size	Unfed	8.0 \pm 0.2	8.5 \pm 0.3	8.3 \pm 0.2	8.3 \pm 0.2
	Fat	8.0 \pm 0.3	7.8 \pm 0.2	8.6 \pm 0.2	8.2 \pm 0.1
	Fat + Vit E	8.5 \pm 0.3	7.7 \pm 0.3	7.6 \pm 0.2	7.9 \pm 0.1
	Annual mean	8.2 \pm 0.2	8.0 \pm 0.2	8.2 \pm 0.1	
Hatching success (%)	Unfed	91.5 \pm 2.3	87.2 \pm 2.6	91.0 \pm 2.6	89.9 \pm 1.4
	Fat	93.1 \pm 2.3	91.7 \pm 2.6	88.0 \pm 2.7	90.6 \pm 1.5
	Fat + Vit E	93.1 \pm 2.0	90.6 \pm 2.3	95.1 \pm 1.5	92.9 \pm 1.2
	Annual mean	92.6 \pm 1.3	89.9 \pm 1.4	91.1 \pm 1.4	

antioxidants such as vitamin E^{22,23}. Daily energy acquisition is a key determinant of winter survival in small passerine species of birds, which can readily fall into negative energy balance when natural food is scarce^{17,24}. However, over-reliance on supplemental energy-rich foods such as fats to the detriment of a balanced diet could be harmful²¹. Furthermore, passerine species cannot store energy to any great extent. Therefore, with the exception of caching species, energy sequestered from the diet in winter is unlikely to fuel later reproduction²⁵. Lipophilic antioxidants, on the other hand, may enhance health through effects on immunity and antioxidant defence^{22,26} and can be stored in large quantities in body tissues^{27,28}. As such, it has been hypothesised that antioxidants in body storage could influence reproduction in later seasons or years²⁹. In addition, the impacts of winter food supplementation on wild bird species are likely to be influenced by stochastic variation in natural food availability^{18,30}. To detect such impacts therefore requires a landscape-scale experimental approach that encompasses multiple sites and years.

Using a three year experimental study conducted across nine woodland sites, we investigated the impacts of winter supplementation of energy- and antioxidant-rich foods on future reproduction in blue tits. Woodland populations of tits were randomly allocated to receive either fat ($n = 3$), fat plus vitamin E ($n = 3$), or remained unfed as a control ($n = 3$). The experiment was conducted over three years (2008–2010), and treatments were rotated amongst sites across years to control for any site-specific effects on breeding performance. Using woodland populations allowed greater control of the context and quantity of food provisioning than could have been achieved within an urban environment. Since natural food and breeding resources are anticipated to be more accessible to woodland birds³¹, our results are predicted to be a conservative estimate of the consequences of winter provisioning within a strictly urban landscape.

Results

There were no significant effects of winter food supplementation on nest box occupancy (binomial GLMM; $n = 467$ out of 1038 boxes occupied over 3 years $\chi^2_2 = 3.17$, $P = 0.21$). The timing of laying and clutch size also did not vary in response to food supplementation, as previously described in Plummer et al. (2013)²¹. Hatching success was marginally higher in fat plus vitamin E-fed populations compared to the other treatment groups (Table 1; binomial GLMM; $n = 362$, $\chi^2_2 = 5.44$, $P = 0.066$). However, compared to unfed controls, parents that had been supplemented in the previous winter produced offspring that weighed less and were structurally smaller, and as a

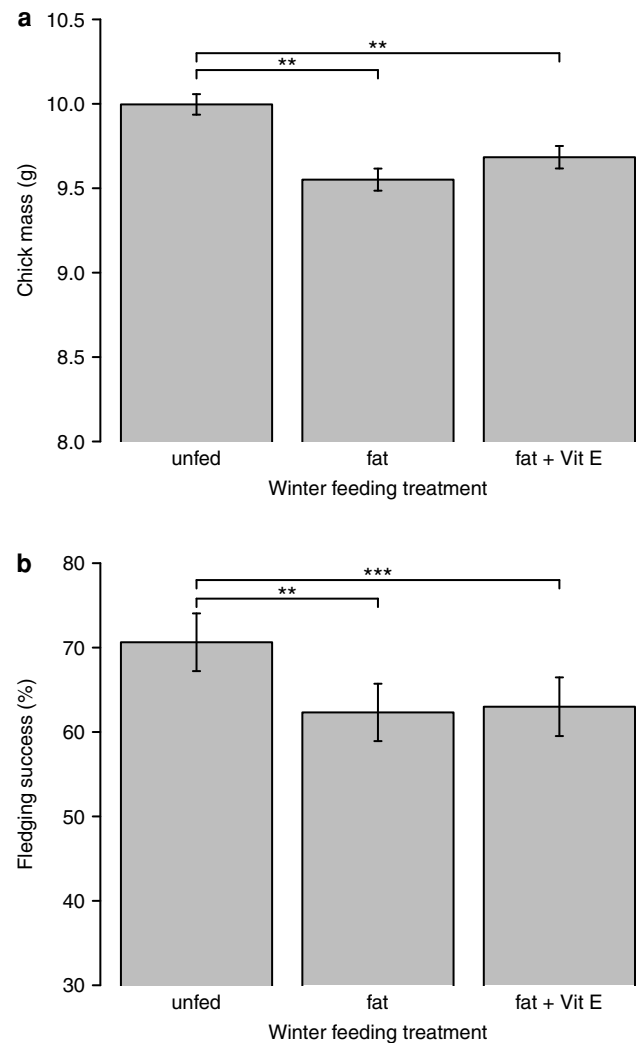


Figure 1 | Differences in (a) chick mass and (b) fledging success in response to winter feeding. Mean \pm s.e.m plotted using raw values. Significance of *post-hoc* pairwise comparisons shown, where ** = $P \leq 0.01$ and *** = $P \leq 0.001$.



Table 2 | Effects of winter food supplementation of parents on (a) chick mass, (b) chick head-bill length and (c) fledging success during the subsequent breeding season

Fixed effect	Factor level	Estimate \pm s.e.m	χ^2	d.f.	P
(a) Chick mass					
Treatment ^a	Fat	-1.749 ± 1.387	10.83	2	0.004**
	Fat + Vit E	-5.033 ± 1.436			
Year ^b	2009	-0.503 ± 0.166	9.01	2	0.011*
	2010	-0.255 ± 0.172			
Hatching date		-0.003 ± 0.021	7.83	1	0.005**
Brood size		-0.240 ± 0.039	35.01	1	<0.001***
Nestling age		0.436 ± 0.093	21.50	1	<0.001***
Treatment \times hatch date ^a	Fat	0.026 ± 0.028	10.52	2	0.005**
	Fat + Vit E	0.093 ± 0.029			
(b) Chick head-bill length					
Treatment ^a	Fat	-0.334 ± 0.101	12.93	2	0.002**
	Fat + Vit E	-0.307 ± 0.103			
Year ^b	2009	-0.550 ± 0.105	8.71	2	0.013*
	2010	-0.245 ± 0.109			
Hatching date		0.018 ± 0.008	4.62	1	0.032*
Brood size		-0.093 ± 0.023	13.32	1	<0.001***
Nestling age		0.549 ± 0.059	74.52	1	<0.001***
(c) Fledging success					
Treatment ^a	Fat	-0.599 ± 0.188	15.21	2	<0.001***
	Fat + Vit E	-0.729 ± 0.194			
Year ^b	2009	0.687 ± 0.179	14.83	2	<0.001***
	2010	0.452 ± 0.194			
Hatching date		0.037 ± 0.018	4.23	1	0.040*

All main effects and significant interaction terms within each minimum adequate model are reported, following stepwise deletion of non-significant terms.
^aWinter feeding treatment relative to unfed controls.
^bRelative to 2008.

consequence fledged an average of 8% fewer offspring (Fig. 1, Table 2). Indeed, the body mass and head-bill length of chicks 12 days post-hatching strongly predicted their subsequent fledging success in all treatments and years (binomial GLMM, $n = 288$; brood mean mass: $\chi^2_1 = 122.97$, $P < 0.001$; brood mean head-bill length: $\chi^2_1 = 70.61$, $P < 0.001$). These deleterious effects were irrespective of the

specific type of food that was supplemented during the previous winter (Fig. 1, Table 2). Furthermore, there was significant annual variation in all breeding measures ($\chi^2 \geq 8.71$, $P \leq 0.013$) except nest box occupancy and hatching success ($P \geq 0.13$). But the effects of treatment were not significantly influenced by year in any instance (treatment \times year interaction: $P \geq 0.096$).

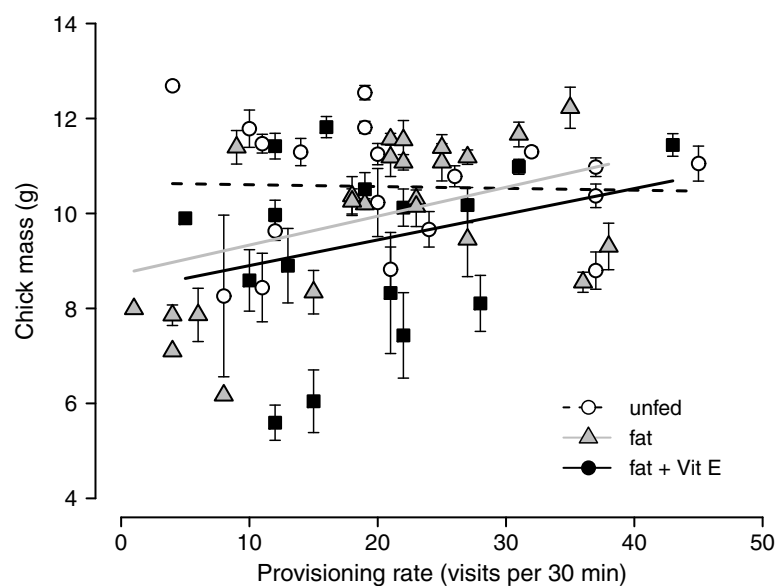


Figure 2 | Relationship between parental nest visit rate and chick mass. Brood means \pm s.e.m are plotted and lines fitted using parameter estimates from the minimum adequate model. *Post-hoc* comparisons showed that the relationship was significantly different for unfed parents compared to fat-fed ($P = 0.046$) and fat plus vitamin E-fed (fat + Vit E) parents ($P = 0.017$).



We predicted that variation in offspring size and survival could result from differences in parental brood-rearing capacity carried-over from the winter feeding treatment. Therefore, we conducted a concurrent investigation to assess the extent to which variation in chick mass at day 12 was explained by parental nest visit rates during the first year of the study. Data were collected using video recordings at 59 nests across all nine woodlands and all three feeding treatments. Parental nest visit rates did not differ significantly amongst treatments (Poisson GLMM; $n = 59$, $\chi^2_2 = 2.05$, $P = 0.36$). However, nest visit rate positively predicted chick mass and this relationship differed amongst treatments (GLMM; $n = 300$, treatment \times nest visit rate interaction: $\chi^2_2 = 8.11$, $P = 0.017$). There was no significant relationship between nest visit rate and the mass of chicks produced by unfed (control) parents, which were heavier on average (Fig. 1a). However, nest visit rate significantly predicted the mass of chicks produced by winter-fed parents of both food supplementation treatments (Fig. 2). Therefore, whilst winter-fed parents did not exhibit a reduction in nest visits *per se*, the quantity and/or nutritional value of delivered food items was clearly diminished.

Discussion

The results of this study cast doubt on the assumption that supplementary feeding of wild birds is invariably beneficial at the population level. Winter food supplementation resulted in reduced breeding success the following spring, several weeks after the provisioning of food had ended. Our results therefore provide an experimental demonstration of a carry-over effect, one of few such examples reported in a wild animal¹⁵. The capacity for within season food supplementation to influence reproduction in wild birds varies amongst sites and across years, for example in response to variation in natural food supply^{18,32–35}. Such variance across years is clearly evident from the results of the present study (see Table 1). Yet, despite taking into account these notable sources of variation by conducting our research at nine woodlands and over three years, the overwhelming pattern which emerged was of deleterious impacts of winter food supplementation on future breeding performance.

Marginal increases in hatching success following vitamin E provisioning suggest that females were able to improve egg investment to some extent following increased antioxidant uptake in winter^{21,36}. However, the negative effects of winter food supplementation on the weight, size and survival of chicks (Fig. 1), and on parental chick-rearing capacity (Fig. 2), suggests that initial investments made by fed parents could not be sustained post-hatching. Although conducted in a woodland environment, the difference in fledging success we report between fed and unfed populations resembles the level of variation in productivity observed between birds in urban and non-urban landscapes. Fledging success is on average 15.4% (95% confidence range: 8.9–21.9%) lower in urban compared to rural habitats for passerine bird species³¹. This difference has been attributed to the multiple costs associated urban living, including increased predation, greater disease transmission, fragmented habitat structure, road collision risk and reduced food availability³¹. Our results support the suggestion that supplementary feeding in urban gardens may also be a contributory factor^{14,31}.

There are at least three possible explanations for our findings. First, winter feeding could have resulted in an unbalanced diet if birds heavily utilised supplementary food, and subsequently entered the breeding season in relatively poor nutritional condition. We are aware of only two previous studies which have experimentally investigated the effects of winter food supplementation on any aspect of future reproductive performance in birds. We recently reported that winter feeding with fat resulted in the production of eggs with relatively small yolks and reduced levels of carotenoid pigments early in the laying season, compared to birds supplemented with fat plus vitamin E or which were unfed²¹. In contrast, Robb et al.²⁰ reported that winter feeding with peanuts led to earlier laying and increased

fledging success in woodland populations of blue tits in a single year of study. One possible explanation for this difference is that the impacts of food supplementation on reproduction depend on the specific nutritional profile of foods. For example, sulphur-containing amino acids are potentially limiting for reproduction in birds, including tits³⁷, although peanuts are a relatively poor source of these nutrients³⁸. Dependency on supplemented fats, on the other hand, while being an excellent source of energy could result in an unbalanced diet with negative consequences for reproduction. We found that reproduction was impaired in populations that had been provisioned with fat, but also in populations which had received fat together with vitamin E – a potent antioxidant and immunostimulant^{22,26}. Therefore, negative effects of a diet rich in fats might override any benefits of antioxidant intake, although this possibility awaits study.

Second, through effects on survival and recruitment, winter food supplementation could have altered the phenotypic structure of breeding populations, such that they included increased numbers of individuals that had relatively low reproductive capacity. Third, winter food supplementation could have caused an ecological trap by encouraging birds to make unsustainable investments in reproduction, at localities which had insufficient natural food availability in spring^{10,16}. However, it is difficult to separate these possible mechanisms using breeding parameters at the population level as presented here. For example, we found no significant effect of winter food supplementation on nest box occupancy, as might be expected if breeding population density increased as a result of winter feeding either through greater survival or attraction to feeders. Although the abundance and utility of natural nesting sites is unknown and as such a true measure of breeding density is unavailable. To disentangle such carry-over effects and better understand the efficacy of bird feeding these possible mechanisms would ultimately require known individuals to be followed through winter and spring, which is almost impossible to do in the wild³⁹.

In conclusion, this study has shown for the first time that winter food supplementation had deleterious consequences for future fledging success. Nevertheless, despite the evidence presented here, it would be premature to conclude that supplementary feeding is inevitably deleterious for wild birds. It would be particularly valuable to assess whether year-round food supplementation can alleviate negative effects of season-specific feeding. For example in the US, Florida scrub-jays living in suburban environments are believed to gain reproductive benefits due to the abundance and predictability of anthropogenic food sources⁴⁰. Furthermore, how we judge the efficacy of food supplementation to wild birds must depend in part on whether impacts are assessed at the level of individuals or populations. For some individuals, the receipt of supplementary food may improve their lifetime reproductive success if they would otherwise have been unable to survive and recruit, or to reach a condition threshold necessary to breed. However, recruitment of such individuals might result in reduced mean productivity at the population level. Whether prolonged winter provisioning could bring about declines in the population sizes of wild birds in the long-term is an important question for future study. We note that there is no indication that overall numbers of blue tits in the UK have declined; in fact numbers have remained stable since 1995⁴¹. However, potential relationships between winter food supplementation and trends in population sizes of birds have not been investigated for any species. With garden bird feeding promoted as a method for conserving declining wild bird populations, these new insights suggest much more needs to be done to fully understand its impacts.

Methods

Study sites and experimental design. The study was conducted at nine deciduous woodlands in Cornwall, UK. Study woodlands averaged 10.7 hectares (± 1.1 s.e.m.) and were situated at least 2 km apart. The predominant tree species were oak (*Quercus* spp.), beech (*Fagus sylvatica*) and sweet chestnut (*Castanea sativa*).



Woodlands were grouped into triplets according to their tree, understorey and ground cover species composition. Within sites, feeders and nest boxes were positioned at an equal density and distribution of *ca.* one feeder and four nest boxes per hectare. Given that woodlands were of a comparable size with nest boxes provided at equal densities, it is unlikely that our findings were influenced by density-dependent effects. The winter food supplementation experiment was conducted over three years from 2007 to 2009 (14 December – 4 March 2007/08; 18 November – 11 March 2008/09 and 2009/10), and potential carry-over effects on reproductive success were investigated during the subsequent springs, 2008–2010. In the first year of the study, each site within a triplet was randomly allocated to a feeding treatment: unfed (control), fat, or fat plus vitamin E. Treatments were rotated within triplets across years, so that every site received all three treatments over the course of the study. Therefore the potential confounding effect of variation in site quality was controlled for in the experimental design.

Winter food supplementation. All feeding stations were provisioned with a new 150 g fat ball every 10 days. Fat balls were produced 1–2 days in advance of provisioning, using solid vegetable fat (Crisp 'n Dry, Princes Ltd., Liverpool, UK) and a small amount of yellow food colouring to increase food uptake by birds (0.125 mL 100 g⁻¹ fat; ASDA Natural Food Colouring, Asda Stores Ltd., Leeds). Fat balls for the fat plus vitamin E treatment group were supplemented with α -tocopherol (T3251; Sigma-Aldrich, Dorset, UK) at a concentration of 100 mg kg⁻¹ fat, a level equivalent to that found in peanuts⁴². Lipophilic antioxidants cannot be provisioned to birds without the use of fat or oil as a 'carrier', and such antioxidants are always co-acquired with lipids in natural foods⁴³. All supplements were prepared using standardised methods²¹, and were predominantly utilised by the target species. Winter misting netting and stable isotope analysis confirmed the use of winter supplemented food by breeding birds.

Breeding success parameters. Nest boxes were inspected every 1–3 days from April to June to determine laying date, clutch size, brood size and fledging numbers. Hatching success was defined as the proportion of a clutch that hatched, and fledging success as the proportion of hatchlings that fledged. Morphometric measures were recorded for all surviving chicks at 12 days post-hatching (± 1 day; mean chick age \pm s.e.m.: 12.19 \pm 0.02, n = 1706, brood n = 319). Body mass (± 0.1 g) was recorded using an electronic balance and head – bill length (± 0.05 mm) measured twice with dial callipers then averaged⁴⁴.

Parental visit rates. Parental visit rates to nests were recorded on day 11 (± 1 day; n = 59) of the nestling phase during the first breeding season (2008). Nest boxes were filmed for 30 min using a Sony DCR – HC37E Handycam at 0630 GMT or 0830 GMT. Chick age and filming time did not differ significantly between treatment groups (GLMs: $P > 0.15$). Parental visit rate, defined as the number of visits per 30 min, was extracted from video playbacks. The sex of parents could not be distinguished from the video recordings, and therefore nest mean visit rates were calculated.

Statistical analyses. All statistical analyses were conducted using R version 2.12.2⁴⁵. Data were analysed using general/generalised mixed effects models (GLMMs) and backwards stepwise deletions of non-significant terms, with alpha set at 0.05. Hatching and fledging success data were fitted using binomial errors and parental nest visit rates using Poisson errors, after checking for overdispersion. Remaining analyses were completed using a Gaussian error distribution, and the normality and homoscedasticity of residuals were checked prior to model simplification. For analysis of nest box occupancy, lay date, clutch size, hatching and fledging success, nest box identity nested within woodland site was specified as the random term to control for pseudoreplication. Brood identity was included as a random term, nested within nest box identity, in analyses of chick phenotypes to control for non-independence of chicks within broods. Nest box identity was excluded from analyses involving parental nest visit rate, as boxes were only used once within a given year. Winter food supplementation treatment, year and lay/hatching date were included as explanatory variables in all relevant models. Brood size at hatching and chick age at sampling were additionally controlled for in chick analyses. All two-way interactions involving treatment were fitted in all models. For *post-hoc* testing, ANOVA was used to compare the minimum GLMM model with replicate models in which two treatment groups under comparison were paired. A total of 467 boxes were nested in by blue tits over the study period. Lay date analysis included all clutches, but for all subsequent analyses clutches with laying breaks > 2 days (n = 23) and/or abandoned prior to incubation (n = 32) were excluded. Only hatched clutches were considered in hatching and fledging success analyses (n = 362). Broods abandoned within one day of sampling activities, or which could not be regularly monitored due to time constraints (n = 28) were excluded from chick size and fledging analyses. These data filters were not biased amongst treatment groups ($P > 0.11$).

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Author contributions

K.E.P., S.B., D.I.L., D.E.C. and J.D.B. designed the research. K.E.P. and J.D.B. wrote the first draft, and the manuscript was reviewed by all the authors. K.E.P. collected and analysed the data.

Additional information

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